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**Running paths to nowhere: Repetition of routes shows how navigating ants
modulate online the weights accorded to cues**

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Abstract

Ants are expert navigators, keeping track of the vector to home as they travel, through path integration, and using terrestrial panoramas in view-based navigation. Although insect learning has been much studied, the learning processes in navigation have not received much attention. Here we investigate in desert ants (*Melophorus bagoti*) the effects of repeating a well-travelled and familiar route segment without success. We find that re-running a homeward route without entering the nest impacted subsequent trips. Over trips, ants showed more meandering from side to side and more scanning behaviour, in which the ant stopped and turned, rotating to a range of directions. In repeatedly re-running their familiar route, ants eventually gave up heading in the nestward direction as defined by visual cues and turned to walk in the opposite direction. Further manipulations showed that the extent and rate of this path degradation depend on 1) the length of the vector accumulated in the direction opposite to the food-to-nest direction, 2) the specific visual experience of the repeated segment of the route that the ants were forced to re-run and 3) the visual panorama: paths are more degraded in an open panorama, compared with a visually cluttered scene. The results show that ants dynamically modulate the weighting given to route memories, and that fits well with recent models suggesting that the mushroom bodies provide a substrate for the reinforcement learning of views for navigation.

Key words: desert ant, view-based navigation, path integration, mushroom bodies, cue weighting

46 **Introduction**

47 **Sisyphus in the desert**

48 Imagine traversing a familiar route home in the blazing red desert of Central
49 Australia, but when the front door looms and the comforts of home beckon, you
50 find yourself inexplicably back where you started your journey. How would you
51 retrace your route? This is the nightmare that we foisted repeatedly on red honey
52 ants, *Melophorus bagoti*, expert navigators wielding and combining multiple
53 strategies (Cheng 2012; Cheng et al. 2009; Cheng et al. 2014; Kohler and Wehner
54 2005). This conundrum for the ants, facing the displacements we call *rewinding*,
55 revealed in detail what happens when their favourite strategy, following well-
56 travelled routes based on the visual panorama, no longer reaps its usual reward of
57 getting home. We were inspired to expand on two accounts on this theme on
58 formicine cousins of *M. bagoti*, North African *Cataglyphis* ants (Andel and Wehner
59 2004; Collett 2014). We here depict in detail how the tiny brain of a desert ant
60 adjusts its navigational toolkit when faced with the drama inflicted by rewinding, a
61 burden for the heat loving red honey ants (Christian and Morton 1992; Muser et al.
62 2005) reminiscent of the task for the mythical Sisyphus, who had to push a rock up
63 a hill only for it to roll down to the bottom again.

64

65 Individually foraging desert ants boast three major strategies for navigating
66 robustly over long distances (Collett et al. 2013; Knaden and Graham 2016;
67 Pritchard and Healy 2017; Wehner 2003): path integration (PI), the ability to keep
68 track of the straight-line distance and direction from the starting point of travel
69 (Collett and Collett 2000; Müller and Wehner 1988; Ronacher 2008; Wehner and

Srinivasan 2003), view-based guidance relying on terrestrial visual (hereafter visual) information learnt from panoramic scenes (Collett et al. 2007; Graham and Cheng 2009; Pritchard and Healy 2017), and systematic search (Schultheiss et al. 2015; Wehner and Srinivasan 1981) to compensate for errors in navigation. Ants combine information from different strategies simultaneously and flexibly (Collett 2012; Legge et al. 2014; Narendra 2007; review: Wehner et al. 2016), perhaps even weighting the cues optimally based on their reliability (Hoinville and Wehner 2018; Legge et al. 2014; Wystrach et al. 2015). Thus, when walking in uncertain conditions, desert ants zig-zag more from side to side, called meandering, and stop and scan the environment more, showing searching mixed with directed navigation (Wehner et al. 2016; Wystrach et al. 2014; Wystrach et al. 2011b). We asked how red honey ants adjust their navigational strategies when their familiar route home does not lead to their entering their abode.

Rewinding to reveal intricacies of running routes without success

Our rewinding technique was inspired by two studies tapping this theme. Andel and Wehner (2004) rewound *C. bicolor* ants as they ran home along a narrow channel decked with landmarks. In their experiment, the ants reached their nest on each trip, only to be dragged out again to re-run the channel. This was as if Sisyphus succeeded in placing the rock at the top of the hill, but then was asked to push a second rock. Rewinding built up the vector calculated by PI. When the ants were tested in a long channel without adorning landmarks—thus without the familiar visual cues defining their route—the rewound ants dashed off in the opposite, nest-to-start-point direction. Collett (2014) allowed individual North African *C. fortis* ants to develop routes across an open environment dominated by a

single conspicuous black cylinder. Collett (2014) then rewound the ants, now with their PI vector indicating zero, just once. Interestingly, some ants appeared confused for a prolonged period before eventually commencing their regular route, suggesting that a single unsuccessful event reduces the trust that ants have in their visually defined direction (Graham and Mangan 2015).

We rewound red honey ants repeatedly without letting them enter their nest. We documented their meandering, scanning, and U-turning to move in the opposite direction. We predicted increases in all these behaviours as rewinding stamps its mark over repeated re-runs. A series of manipulations let us delve into three factors contributing to reduced confidence in route running: the length of the vector in the opposite, nest-to-feeder direction built up by rewinding, the memories of the particular visual cues along the rewind portion of a route, and the structure of the visual environment. In addition, we tested whether our experimental manipulations can lead to impacts even following long intervals, to test if long-term route memories are updated because of the rewinding process.

Methods

Animals

Thermophilic, diurnally foraging red honey ants, *Melophorus bagoti*, are found throughout semi-arid Central Australia (Christian and Morton 1992; Muser et al. 2005). Two nests at a field site on a private property ~10 km south of Alice Springs, Australia were used.

Experimental set up

The scenery surrounding the nests consisted of grass tussocks, mostly of the invasive buffel grass (*Cenchrus ciliaris*), bushes, trees, and a few low buildings (Fig. SI1). The test area, however, was cleared of vegetation. One of our experimental nests (Nest 1) had plants near the nest and test area, making for proximal cues in the surrounding panorama, while Nest 2 lay in an open area with little vegetation in the vicinity of the nest.

A square-shaped feeder made of plastic (15 × 15 × 9 cm deep) was sunk into the ground 10 m from a nest. At Nest 1, an enclosure of smooth plywood low walls 10 cm high was constructed around feeder and nest (Figs. 1A, SI1). Both these walls and the walls of the feeder were too slippery for the ants to climb. The feeder was provisioned with mealworm and cookie pieces. During training, sticks were used as bridges, allowing foraging ants to climb out of the feeder.

The enclosure walls were used to form two tracks (Figs. 1A, SI1). A narrower track free of obstacles served as the outbound route for the ants, while the wider track, with obstacles along the way, served as the homebound route. Strings wound around tent pegs were used to make a grid of 1-m squares. The grid

135 allowed experimenters to transcribe the paths of homebound ants onto similarly
136 gridded paper (i.e. copies of Fig. 1A). For a coordinate system, we labelled the nest
137 0,0, while the feeder was labelled 0,-10, with metres as units. Two sections of the
138 homeward path were labelled Section A (from $y = -10$ to $y = -7$) and Section B ($y =$
139 -6 to $y = -3$).

141 **Procedure**

142 All ants were trained to be familiar with the homeward route before being
143 tested. An ant arriving at the feeder for the first time was painted with one dot of
144 enamel paint (Tamiya™ brand), in a colour that denoted the day of arrival at the
145 feeder. Ants were tested after at least two days from the day of their first arrival.
146 Each ant was tested on only one occasion.

147 Ants were tested one at a time. For tests, the sticks were removed from the
148 feeder for the duration of the test. The test ant was captured in a dark tube and
149 released with its food just in front of the feeder or further along the route,
150 depending on the test. To ensure homing motivation, only ants holding on to their
151 piece of food were tested. The test ant was allowed to run home but typically
152 captured in a dark tube along the route or just before it entered the nest to be
153 returned to a release point to run again, a process we called rewinding. Similarly,
154 in order to manipulate the PI state of ants prior to the start of rewinding, ants
155 could be allowed to complete an outward run to the feeder before being captured
156 and returned to the vicinity of the nest for another outward run. Variations on this
157 theme made up the test conditions.

Test conditions

We investigated four questions by making ants repeat routes. **1) What is the effect on habitual route guidance of conflicting path integrator states caused by rewinding?** To do this we took ants that had run to the feeder either once or three times; thus we had groups starting with different PI home vectors and different conflicts between visual guidance and PI at each stage in the rewinding.

2) What is the specific effect of visual repetition from rerunning one portion of the route? Here ants were manipulated so that they only had repeated exposure to a specific portion of the route. This tests if rewinding modulates visual guidance generally, or affects the specific views encountered in the rewound section especially adversely. **3) How persistent are the effects of repeating a route?** By asking whether route repetition influences navigational memories even after long periods, we can ask if rewinding changes long term memories. **4) What is the influence of environmental structure on how route repetition influences routes?** We tackled this question by asking ants to repeat visually simple or visually complex routes and asking if behavioural changes differ according to the type of visual scenery.

Each of these questions was tackled via several test conditions. Each condition was given a number/letter code (Table 1), with a short description. Full details of test conditions are given in the supplementary materials.

Table 1 about here

Data analysis

All recorded test paths were digitised with GraphClick™ for further analysis, delivering a series of coordinates describing ants' paths. We analysed Scan rate, defined as the number of scanning bouts displayed per metre of path travelled (with scans as defined by Wystrach et al., 2014), and Meander, defined as the mean of the angles formed between successive 30cm path segments (following Wystrach et al., 2011b). Most of the analysis consisted of standard parametric statistics, with Condition as a between-subjects factor, and rewinding trial number as the repeated measure. But the trial number on which an ant first performed a U-turn and travelled back in the nest-feeder direction was analysed using non-parametric statistics.

Table 1. Summary of test conditions

Test condition		Key manipulations
1a	One-outbound re-runs	1 outbound trip repeated homebound trips
1b	Three-outbound re-runs	3 outbound trips before arriving at feeder repeated homebound trips
1c	Nine-captures in a single homeward run	captured 9 times on 1 trip home test after 10th capture near nest
2a	Re-run of Section A only	ran Section A 4 times test on 5th trip
2b	Re-run of Section B only	ran Section B 4 times test on 5th trip
3a	Re-run of Section A with delay	ran Section A 4 or 5 times delayed for 24 h before test
3b	Re-run of Section B with delay	ran Section B 4 or 5 times delayed for 24 h before test
3c	Re-run of Sections A and B with nest return	ran Section A 4 times, then went home next foraging trip; ran Section B 4 times tests immediately after running Section B (as zero-vector ant) and on next trip to feeder (as full vector ant)
4a	One-outbound re-run open (Nest 2)	1 outbound trip (Nest 2), open repeated homebound trips
4b	One-outbound re-run with clutter (Nest 2)	1 outbound trip (Nest 2) with landmarks added near route repeated homebound trips

Fig. 1 about here

Results

The effect on visual route guidance of route repetition and conflicting path integrator states

a) Conditions 1a and 1b: 1 or 3 outbound runs before repeating homeward route

To investigate how rewinding and PI state influence path characteristics we allowed ants to have either one (Condition *1a*) or three foodward runs (Condition *1b*) from nest to feeder before being allowed to grab some food and beginning re-runs of the homeward route. Thus, at the start of the rewinding process ants either had a normal PI home vector or had accumulated a vector of three times the magnitude of the normal home vector. Therefore, the group with a single outbound journey experienced a conflict between PI and visual guidance from the 2nd rewinding run while the group with three outbound journeys experienced a conflict from the 4th rewinding run. Following those points in the rewinding sequence, the conflict for both groups increased between the ant's PI system and the direction indicated by visual cues, as the PI home vector got longer in the nest-to-feeder direction. Over 6 runs, both groups of ants became more variable in their paths (Fig. 1B), meandering and scanning more. Our formal measure of Meander was higher in ants that had a greater conflict between visual guidance and PI because they had experienced only one outbound run to the feeder (Condition *1a*) than in ants that had experienced three outbound runs before reaching the feeder (Condition *1b*, Fig. 1C). An analysis of variance found statistical significance in all effects: Condition (one outbound trip vs. three outbound trips, $F_{1,156} = 56.01$, $p < 0.0001$), trials ($F_{5,156} = 14.57$, $p < 0.0001$), and their interaction ($F_{5,156} = 3.42$, $p = 0.006$). Ants also scanned more frequently over repeated runs home (Figs. 1D, F),

and ants with three outbound runs scanned less often than ants with one outbound run (Fig. 1D). An analysis of variance found significant main effects of Condition ($F_{1,156} = 28.35, p < 0.0001$) and trials ($F_{5,156} = 16.07, p < 0.0001$), but not a significant interaction ($F_{5,156} = 1.53, p = 0.184$). Scans were not evenly distributed across the length of the route home (Fig. 1B). Both ants with one outbound run and ants with three outbound runs scanned most in the early part of the route, especially just before the first set of barriers at -7.

We further analysed how ants lost confidence in their visually guided route by looking for the point at which the PI direction finally overrode the visually defined direction and ants U-turned and began travelling in the nest-to-feeder direction. Ants with 3 outbound runs before rewinding had a higher survival curve (proportion of ants that had still not performed a U-turn by a given trial number) than ants with 1 outbound run (Fig. 1E). This difference, however, did not reach statistical significance (non-parametric test on number of trials before the first turn-back: $Z = 1.68, p = 0.092$). After U-turning, ants typically ran for a long distance away from home, that is, in the direction of their negative path integration vector, confirming that ants had indeed accumulated a large vector, as observed by Andel and Wehner (2004).

b) Condition 1c: Nine captures on one trip home

As a control for the repeated capturing and disruption of the homebound journey, we captured ants 9 times on one journey home, resulting in a zero-vector ant being tested after the 10th capture. Repeated capturing had no notable effect

on the ants' navigation (Fig. SI2). The ants performed much like zero-vector ants that had been captured only once near their nest. Ants in Condition *1c* had a closer resemblance to ants with a same PI state (Condition *1a*, 2nd release) than to ants after the same number of captures (Condition *1a and 1b*, 10th release, Supplementary Results SI2). While we cannot rule out that repeated captures may have some effect on paths, the changes in behaviour that arise from repeatedly running the home path must be attributed mostly to the increasing vector length of path integration in the negative (nest-to-feeder) direction and/or repeated viewing of the scenes along the route, which is examined next.

The specificity of the effect of visual repetition on route disturbance

Condition 2a and 2b: Re-running Section A or Section B repeatedly

In these conditions, ants re-ran only Section A or only Section B four times and were then tested on the entire route (Fig. 2A). In comparison with ants that re-ran the entire route repeatedly, their meander and level of scanning were similar. We then examined Scan rates and Meander for the particular route sections (A or B) of the entire route. Ants that had re-run Section A repeatedly scanned and meandered more in Section A compared to ants that had re-run Section B (Figs. 2B, D). On the contrary, on Section B of the route, it is ants that had re-run Section B that displayed a higher Meander and Scan rate (Figs. 2C, E). The statistical analysis is combined with Condition 3 in the next sub-section.

Fig. 2 about here

The persistence of route repetition effects

a) Condition 3a and 3b: Re-running Section A or Section B with delay before testing

As in the two previous conditions, ants in Conditions 3a and 3b re-ran Section A only or Section B only four times (Fig. 2). Ants were then held for 24h before being tested on the entire route. The delay served to test if the disruption caused by repeatedly running a segment persists. Results again show a persistent increase in meander and scans specific to the route section that the ants had repeatedly re-run the day before (Fig. 2). Also, the 24h-delay increased the disruption as compared to ants that had performed the test immediately after the 4 rewinding trials (Fig. 2).

For each of Meander and Scan rate on the 5th (test) run, we combined Conditions 2a, 2b, 3a, and 3b in a 3-way analysis of variance, with Condition (rewound on A or B) and delay (immediate test vs. 24-h delay) as between-subjects factors, and section (A or B) as repeated measure. For Meander, the analysis found significant main effects of all three factors (Condition: $F_{1,97} = 8.64, p = 0.0041$; delay: $F_{1,96} = 41.84, p < 0.0001$; section: $F_{1,96} = 55.16, p < 0.0001$). A significant Condition by section interaction was also found ($F_{1,96} = 70.53, p < 0.0001$), while the other interactions were not significant (Condition by delay: $F_{1,96} = 0.38, p = 0.5383$; delay by section: $F_{1,96} = 2.87, p = 0.0935$; Condition by delay by section: $F_{1,96} = 1.82, p = 0.1803$). For Scan rate, the analysis of variance revealed significant main effects of Condition ($F_{1,96} = 8.89, p = 0.0036$) and delay ($F_{1,96} = 12.61, p = 0.0006$), but not of section ($F_{1,96} = 0.25, p = 0.6167$). Each of the interactions was found to be significant (Condition by delay: $F_{1,96} = 6.18, p =$

0.0147; Condition by section: $F_{1,96} = 36.26, p < 0.0001$; delay by section: $F_{1,96} = 11.77, p = 0.0009$; Condition by delay by section: $F_{1,96} = 8.59, p = 0.0042$).

b) Condition 3c: Section A re-run with return to nest

In this condition, ants re-ran Section A 4 times, went inside the nest, reached the feeder again and then re-ran Section B 4 times, so that on a test, they were in a similar path integration state to ants in Conditions 2a and 2b. Compared with Condition 2a, in which ants did not enter their nest before the test, entering the nest did not affect the ants' behaviour in Section A, whereas having a full vector reduced Meander and Scan rate (Supplementary Results, Fig. SI3).

Influence of the visual environment on route disturbance

Conditions 4a and 4b: Re-running with or without proximal objects (Nest 2)

These two conditions were implemented with Nest 2, which lay in a rather open area. Ants re-ran the route home repeatedly without (open landscape, Condition 4a) or with (cluttered landscape, Condition 4b) added proximal objects around the route (Fig. SI4A). The presence of nearby proximal objects mitigated the detrimental effects of repeated route-running, or, to put it another way, having an open landscape along the route meant the effects of repeated route-running were more severe (Fig. SI4B). Ants performed their first U-turn after fewer re-runs in the open landscape (Condition 4a), and they scanned and meandered a little more, compared with their performance in the cluttered landscape (Fig. 3). For Meander and Scan rate on the first three trials, we conducted a mixed analysis of variance with Condition (open or cluttered landscape) as the between-subjects

factor and trials as repeated measure. For Meander, both main effects reached significance (Condition: $F_{1,81} = 9.84$, $p = 0.0023$; trials: $F_{2,81} = 18.45$, $p < 0.0001$), but the interaction did not reach significance ($F_{2,81} = 2.47$, $p = 0.091$). For Scan rate, only the main effect of trials reached significance ($F_{2,81} = 17.72$, $p < 0.0001$). The main effect of Condition ($F_{1,81} = 1.57$, $p = 0.214$) and the interaction ($F_{2,81} = 0.46$, $p = 0.636$) did not reach significance. For the trial on which ants first turned back, the difference between conditions was significant by a non-parametric test ($Z = 3.13$, $p = 0.0018$).

Fig. 3 about here

Discussion

Repeated rewinding produces a Sisyphean task for the ants, in which following their route fails to land them in their nest. The red honey ants transformed their route running, meandering and scanning more and eventually giving up and U-turning in the opposite, nest-to-feeder direction, following the commands of path integration. We built on earlier results on this theme (Andel and Wehner 2004; Collett 2014) to elucidate factors that contribute to the ants' reduced confidence in the familiar visually defined route, normally a favoured strategy of experienced *M. bagoti* foragers (Cheng et al. 2009; Cheng et al. 2014; Kohler and Wehner 2005; Wystrach et al. 2011b).

Factors affecting route-following

As the first factor, increasing conflict between visually guided directions and path integration leads to path degradation. Given an equivalent number of route repetitions, ants with a greater conflicting PI vector length showed more

meandering and pausing to scan, and eventually performed route U-turns earlier in the rewinding sequence (Fig. 1). This shows that learnt visual guidance and PI are simultaneously active in influencing behaviour even when in 180° directional conflict (as observed in Collett 2012; Freas and Cheng 2017; Legge et al. 2014; review: Wehner et al. 2016). Additionally, it shows that the weight given to PI increases with vector length (as shown in Wystrach et al. 2015).

As a second factor, viewing a portion of a route without getting home reduces confidence in that particular segment of the route. Seeing familiar scenes on the route without getting home combines elements of extinction and aversive conditioning of the visually-defined route, without being fully analogous to either learning phenomenon. After repeating a section of a familiar route over and over again, and even in the absence of a strongly conflicting PI vector, ants displayed strong disorientation specifically on that section (Fig. 2). Interestingly, this specific effect is still evident in the routes of ants even after a 24h delay, and whether or not ants had been allowed inside their nest (Figs. 2, SI3). The fluent route-following behaviour did not recover spontaneously, that is, after the mere passage of time without further relevant training. The trust in route-following also did not recover after a single success (see SI3). These results imply that the route recapitulations change long-term memory, rather than causing short-term adaptation. They also illustrate the dynamics of continuous learning in a naturalistic task, highlighting this study's brand of experimental ethology of learning (Freas et al. 2019).

As the third factor, the course of route changes with rewinding depends on the structure of the visual environment. This was suggested by Collett (2014; see

also Graham and Mangan 2015) to explain why path decrements from repetition are not observed in all experimental situations. Here we showed that when ants were navigating in an open environment, as opposed to the same environment with additional proximal visual clutter, path disruption arose after fewer repeated runs, and thus with a weaker conflict between visual guidance and PI (Figs. 3, SI4). It could be that because aversive conditioning sets in for specific views experienced during a familiar route home, the slowly changing visual panorama along an open route provides a larger and longer window to modify the valence of specific views.

Adaptive use of information based on reliability

Ants adjust the weight given to different navigational systems based on many factors. For path integration, more weight is assigned to longer vectors (Hoinville and Wehner 2018; Merkle et al. 2006; Merkle and Wehner 2010; Stone et al. 2017; Wystrach et al. 2015) and for visual guidance, more weight is accorded to familiar views (Legge et al. 2014), unambiguous views (Huber and Knaden 2017), and more recently encountered views (Freas and Cheng 2017). All these rules of thumb make intuitive sense. Here we show that the weight attributed to view-based route guidance also depends on experiences of failure, which reduce the weight given to visually guided route segments. This adjustment does not depend on immediately available information, but requires the accumulation of information over time. That is, this second-order information about the reliability of a navigational system requires a form of memory.

Ants also have reduced trust in their visual route memories when travelling through open environments. Functionally this makes sense as the perceived change in scenery while moving depends on the proximity of the surrounding objects. One can estimate one's position based on memorised views more accurately and precisely if the environment is cluttered (Schultheiss et al. 2013; Zeil et al. 2003). Thus, provided that the current view is equally familiar, visual guidance should be trusted more in cluttered than in open environments. Taken together, our results highlight nuances in the way that ants weight their navigational tools, opening up questions regarding the mechanistic basis of navigation.

The neural basis of flexible route guidance

Our findings give firm support for the role of associative learning in view-based navigation. According to current literature, the mushroom bodies (MB) undergird associative learning in insects (Aso et al. 2014; Bazhenov et al. 2013; Cohn et al. 2015; Galizia 2014; Peng and Chittka 2017; Perry et al. 2013; Webb and Wystrach 2016), including the learning of views that can guide familiar routes (Ardin et al. 2016; Cruse and Wehner 2011; Hoinville and Wehner 2018; Webb and Wystrach 2016). MB support reinforcement learning by separating patterns in the input, and assigning positive valences to positively reinforced patterns. In travelling routes, ants move forward when experiencing familiar reinforced views, and turn more or scan when the view is not similar to reinforced views (Kodzhabashev and Mangan 2015; Lent et al. 2010; Wystrach et al. 2011a; Zeil et al. 2014). We think that rewinding reduces the valence associated with views, also

making ants increase meandering and scanning. Functionally, these behaviours expose the navigator to novel views of the environment, allowing ants to explore new options for reaching home. In general, online updating of the valence of visual route memories could be adaptive in natural foraging. For instance, aversive associations with views might help ants to learn to detour around an obstacle or a trap, a form of avoidance learning (work in preparation).

Like most experimental treatments, our manipulations do not reflect conditions experienced naturally by ants. But the manipulations enabled us to explore the mechanisms underlying natural navigation. We found support in the foraging of ants in their natural habitat for a general point regarding information processing: in general, if a cue is perceived repeatedly and independently of a reward, it is not a good predictor of that reward, and the weight attributed to it should be lowered. In addition, for navigation, a cue perceived repeatedly is not a good indicator of one's current position, and should be ignored. Using our rewinding method enabled us to clearly show that ants do indeed apply this principle, and start to reduce their confidence in views if these are no longer reliable.

Conclusions

The rewinding method asks ants to repeat portions of a familiar route. By implementing rewinding, we have shown dynamic online processing in cue integration, with ants incorporating all route experiences to update route memories and cue weightings. For cue integration, ants use more than heuristics based on immediately available information, such as the current PI vector length

or the current visual familiarity. They also adjust the weight attributed to route memories based on information accumulated over successive trials, that is, over time. Moreover, their fine-tuning depends on how informative the visual scenery is, as reflected in the number of nearby objects (cluttered vs. open environments). We think that reinforcement learning is central to all these experience-dependent modifications. Further investigation of these dynamic learning processes, in conjunction with modelling the associative learning in navigation implemented in the mushroom bodies of insects, will be particularly informative.

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Competing interests

The authors declare that they have no competing interests.

Ethics

Australia has no ethical regulations regarding work with insects. The experimentation carried out was, moreover, non-invasive, and no long-term aversive effects on the nests or on the tested individuals were observed.

Author contributions

AW, SS, and PG conceived and designed the study. AW and SS carried out experimentation and digitised the paths. AW analysed the data. AW drew the figures. PG and KC drafted the manuscript. All authors revised the manuscript and agreed on the final version of it.

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Data accessibility

The raw data files and a file containing explanations are in the supplementary materials.

Supplementary information

Supplementary materials for this article are available at

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598 **Fig. 1. Paths are more disturbed following repetition and cue conflict. A.**
 599 Schematic of the nest and feeder surrounds. A one-way system meant ants had a
 600 clear outward journey before being captured at the feeder. The return journey
 601 included two sections of interest: Section A, which was relatively clear of objects,
 602 and Section B, which was visually cluttered. Diagonal lines represent baffles that
 603 were used to make a clear distinction between Section A and B and also to reduce
 604 wall following. Replicas of this schematic were used to transcribe the routes of ants
 605 during tests. **B.** Paths of ants for their first 6 re-runs of the homeward path. Upper:
 606 Ants had performed only one outward run before the start of the rewinding.
 607 Lower: Ants with three outward runs before rewinding. The locations of bouts of
 608 scanning were also recorded. The spatial distribution of scans for both conditions
 609 are plotted, with dark shades representing an increasing probability that scans
 610 would have been observed within a 0.25m² grid square. **C and D.** For the
 611 conditions with one and three outbound runs, Meander and Scan rate are
 612 compared, respectively. Only ants with at least six trials without U-turn were used
 613 in this comparison. Plots show means with standard deviations. Because of the
 614 different numbers of outbound runs, ants experience a zero-vector state (ZV) on
 615 different run numbers. For additional reference the curve of mean Meander and
 616 Scan rate for the three-outbound group is also shown offset by two runs to allow a
 617 comparison adjusted for PI state (dashed line). **E.** Survival curves for ants in the
 618 one- and three-outbound groups. Lines show proportion of ants still remaining,
 619 with ants removed from the groups once they have performed a U-turn on a
 620 homeward run. **F.** For the one-outbound (upper) and three-outbound (lower)
 621 groups histograms show the change in the number of scanning bouts from one run

to the next. The positive skew shows that scans become more frequent with increasing number of re-runs

Fig. 2. Ants show more disturbance in the sections in which they were rewound. After arriving at the feeder on their first trip ants were allowed to run the first part of the route (Section A) before being rewound to the feeder position a total of four times, or they were taken from the feeder, with their food, to the mid-point of the route and allowed to complete the Section B before being rewound to the mid-point (again for four trips). Thus we have two groups of ants with rewinding experience in Section A or Section B only. **A.** Paths of ants given rewinding in Section A (top) or Section B (bottom). For purposes of clarity, rewinding trials show paths that were selected randomly from a larger number of paths. **B,C.** Meander of ants in Section A (B) or Section B (C). **D,E.** Scan rates of ants in Section A (D) or Section B (E). The box plots show the median (middle line in the box), the 25th and 75th quartiles (bottom and top of the box, respectively), and 1.5 times the interquartile range (whiskers), as well as outliers (individuals beyond the range of the whiskers)

Fig. 3. Ants show more path disturbance after rewinding in an open landscape vs. a cluttered landscape. In order to investigate the impact of visual clutter on route changes, we repeated the basic rewinding protocol (see Fig. 1) with two groups of ants at a second nest site. The nest environs were left open or had visual clutter added. **A.** The proportion of ants that have not turned back in the open and cluttered landscape of Nest 2, and in the landscape of Nest 1, which was

646 visually cluttered (data from Fig. 1E). **B.** Meander in the paths of ants rewound in
647 the open or cluttered landscape of nest 2 (mean \pm s.d.). **C.** Scan rate in the ants
648 rewound in the open or cluttered landscape of nest 2 (mean \pm s.d.)
649

Fig. 1

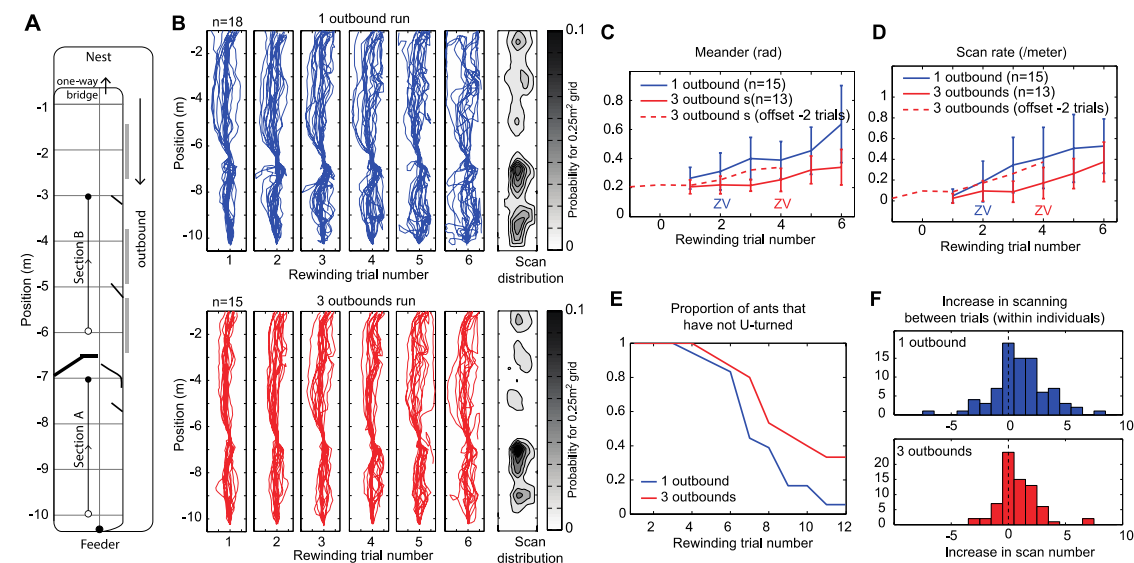


Fig. 2

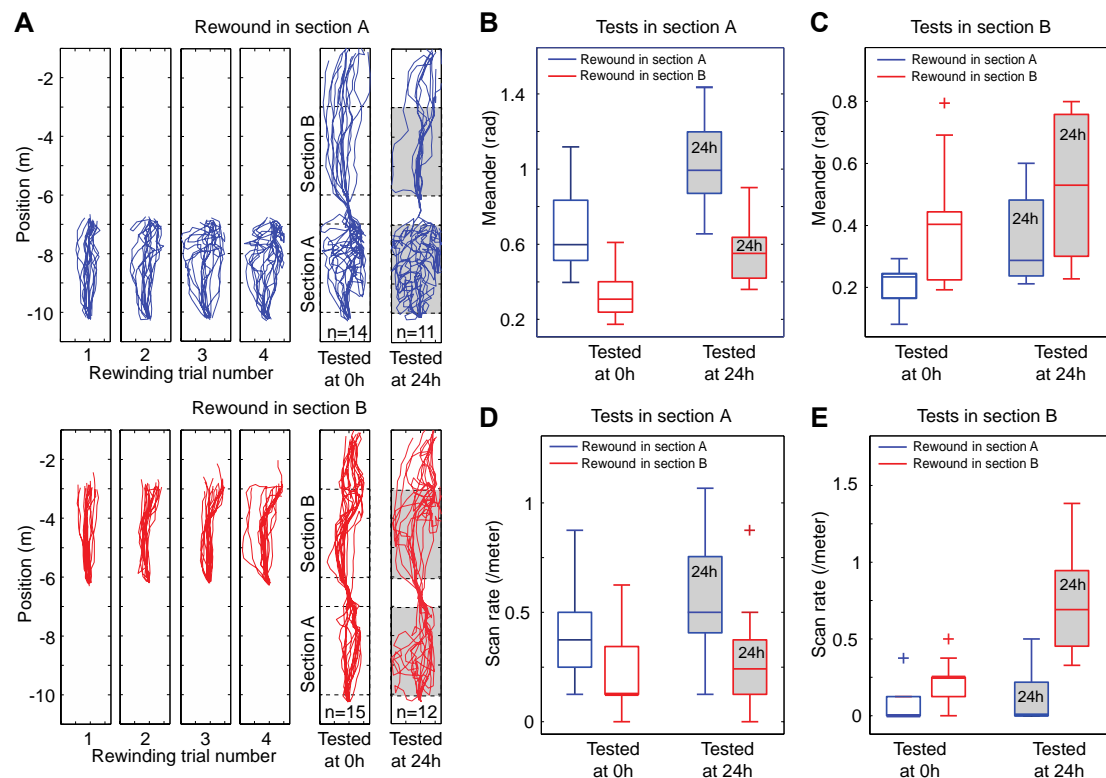
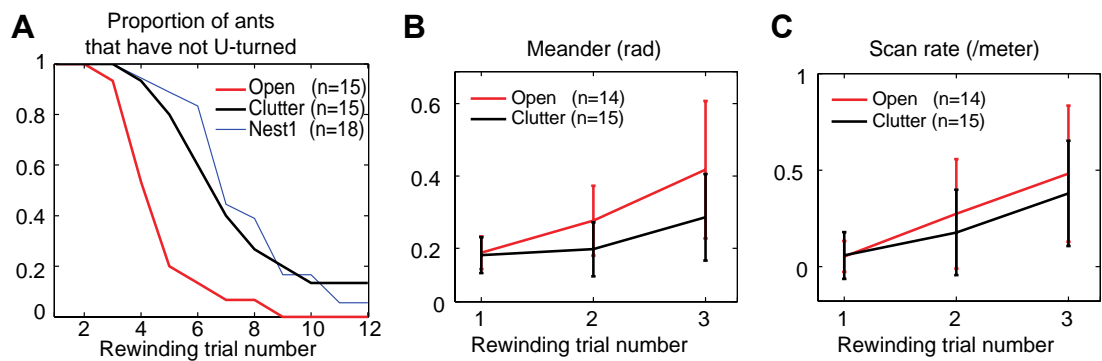


Fig. 3



Supplementary materials

Running paths to nowhere: Repetition of routes shows how navigating ants modulate online the weights accorded to cues

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Supplementary methods

Detailed procedures of each condition

1) The effect on visual route guidance of a conflicting path integrator state

1a. One outbound trip before re-runs. A trained forager was allowed to run home with a piece of food. It was captured just before entering the nest and re-released just in front of the feeder. On the next trip home, the process was repeated. Ants continued to re-run the route home until they U-turned during a homeward route or completed a maximum of 16 runs. U-turns were defined as moving in the nest-to-feeder direction for at least 1 m.

1b. Three outbound trips before re-runs. A trained forager from Nest 1 arriving at the feeder was captured before she grabbed any food. The ant was transported back to her nest and released just in front of the nest. Such a motivated forager then re-ran the route to the feeder, where it was captured once more before seizing any food. It was transported back to the nest once again, and released to run to the feeder for a third time. These ants had thus run the outbound route 3 times, and accumulated a PI home-vector of ~30 m. The ant was then treated in the same manner as Condition *1a*.

1c. Nine captures in a single homeward run. A trained forager dropping into the feeder was allowed to seize a bit of food. The ant was released to run home, and captured at 0,-9 and held in the tube for 20 s. It was released again at 0,-9, and captured again at 0,-8. After being held for 20 s, it was released again at 0,-8. Every metre of travel on the route back, the returning forager was captured, held for 20 s, and released again at the location where she was captured. At 0,-1, the test ant had been captured and released 9 times en route home. Just before the nest, it was captured once more, this time returned to the feeder (0,-10) for one last run home. On this run home, the ant had

run off the 10 m accumulated on the outbound trip, and had no home vector from path integration. In the literature, these ants are called zero-vector (ZV) ants. This condition allowed us to examine the effect of repeated captures without the large accumulation of a path integration home vector pointing in the nest-to-feeder direction.

2) The specificity of visual repetition on route disturbance

2a. Re-run of Section A only. A trained forager arriving at the feeder was allowed to grab a piece of food. The ant was released to run home, and captured at $y = -7$, the end of Section A, and returned to the start. Ants on this test ran Section A 4 times, and then were allowed to run the whole route home on the 5th trip.

2b. Re-run of Section B only. A trained forager arriving at the feeder was allowed to seize a bit of food. The ant was released to run home at 0, -6, the start of Section B, and captured at $y = -3$, the end of Section B, and returned to the start of Section B. Ants on this test ran Section B 4 times, and then were released at the beginning of Section A and allowed to run the whole route home (Section A then B) on the 5th trip.

3) The persistence of effects from route repetition

3a. Re-run of Section A with delay. In this test condition, a forager with food re-ran Section A 4 or 5 times in the same manner as Condition 2a. On the ant's last run, however, it was captured and placed in a dark tube for 24 h. A piece of cotton wet with honey water was provided in the tube. After the delay, the ant was released with food in front of the feeder to run the whole route home once.

3b. Re-run of Section B with delay. In this test condition, a forager with food re-ran Section B 4 or 5 times in the same manner as Condition 2b. Paralleling Condition 3a, on the final run of Section B, the ant was captured and held for 24 h. After the delay, the

ants in this condition were also released with food in front of the feeder to run home once.

3c. Re-run of Sections A and B with nest return. The forager ran Section A 4 times in the manner of Condition 2a; she was then captured and painted with additional colour for individual identification. The test ant was released with its food at the nest, and was allowed to enter the nest. On the forager's next trip to the feeder, she was captured and treated in the manner of Condition 2b: the ant ran Section B 4 times to reach the same PI state as groups 2a and 2b and was released in front of the feeder and recorded along Section A. For any ants that returned after this test, it was tested as a full-vector (FV) ant by being released immediately at the feeder.

4) The influence of environmental structure on effects of route repetition

These conditions were conducted on Nest 2, either within a rather open scene or with additional proximal objects added.

4a. One outbound trip before re-runs in open environment (Nest 2). This test condition was conducted on ants from Nest 2, which had an open area around the nest, replicating the manipulations used in Condition 1a with Nest 1.

4b. One outbound trip before re-runs in cluttered environment (Nest 2). This condition was also conducted on ants from Nest 2. It repeated the manipulations of Condition 4a, except that experimentally provided objects were added in the vicinity of the nest to add visual clutter. Black cylinders, some of them tall, and three rectilinear bamboo baskets were strewn on the sides of the route home.

Fig. S11. Photo of test set up at nest 1.



Fig. S11. Photo of test set up at nest 1. The enclosure walls were used to form two tracks. A narrower track free of obstacles served as the outbound route for the ants, while the wider track, with obstacles along the way, served as the homebound route. The ant's nest entrance opened onto the outbound track, forcing the outbound ants to travel on that track, at the end of which ants could drop into the feeder. The stick bridge that ants used to exit the feeder then forced ants onto the homeward track, at the end of which a one-way ramp allowed ants to drop into the nest. The roof of a tent-like pagoda stood over the central part of the track and plywood baffles (~120 cm wide and ~120 cm tall) were used to segment the route and reduce wall-following behaviour. Strings wound around tent pegs were used to make a grid of 1-m squares.

Supplementary Results

SI2. Being captured 10 times during one trip home does not disrupt route following

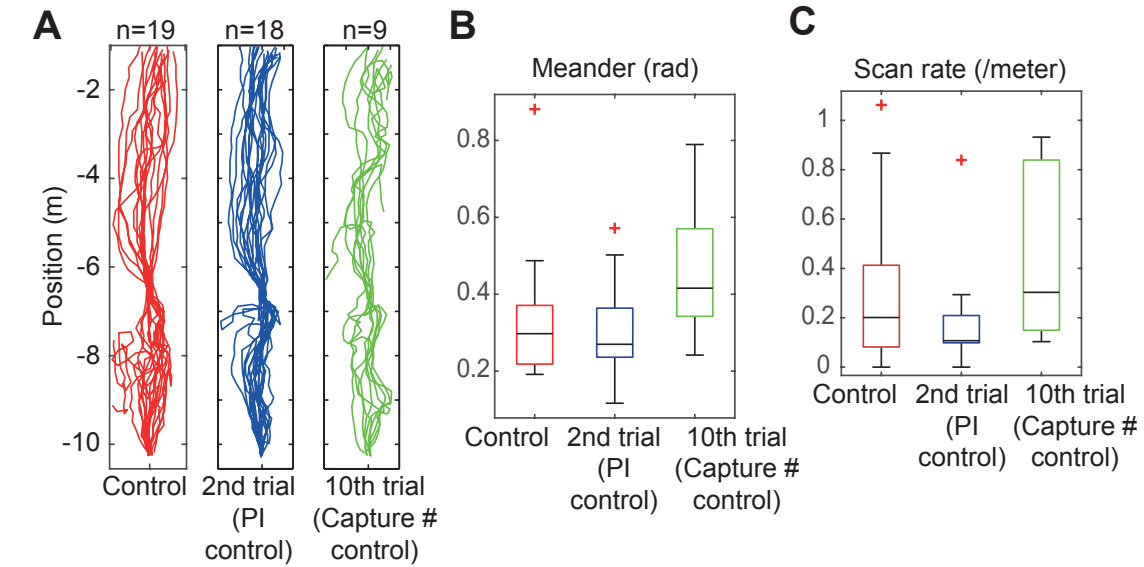


Fig. SI2. Being captured 10 times during one trip home does not disrupt route following. **A.** Paths from the control group captured 10 times during one trip home (control), compared with ants that were captured after completing the full distance to the nest on their second trip and on their 10th trip in Condition 1a. **B.** Meander in the control group captured 10 times during one trip home (control), compared with ants that were captured after completing the full distance to the nest on their second trip and on their 10th trip in Condition 1a. **C.** Scan rate in the control group captured 10 times during one trip home (control), compared with ants that were captured after completing the full distance to the nest on their second trip and on their 10th trip in Condition 1a. The box plots show the median (middle line), 25th and 75th percentiles (bottom and top of box), and 1.5 times the quartile (whiskers), and outliers beyond that range. On the second trial, the ants were equivalent to the controls in PI, while on the 10th trial, the ants were equivalent to the controls in the number of times they had been captured.

SI2 Results

We calculated an evidence ratio to investigate whether ants in Condition *1c* (control in Fig. SI2) had a closer resemblance to ants with a same PI state (Condition *1a*, 2nd release) or to ants after the same number of captures (Condition *1a and 1b*, 10th release). The evidence ratio comes from Bayesian statistics comparing the support of the data from Condition *1c* for one vs. the other hypothesis. For each hypothesis, a non-parametric test comparing Condition *1c* with another condition generated a Z score. The y value of each Z score ($f(Z)$) served as an estimate of the evidence for that hypothesis; the ratios of these y values were computed. For both Meander and Scan rate, ants in Condition *1c* resembled more closely ants with the same PI state (Meander: evidence ratio 12.93; Scan rate: evidence ratio 1.68). These ratios (both being above 1) mean strong evidence in the case of Meander and anecdotal evidence in the case of number of scans. These analyses suggest that the changes in behaviour from rewinding are mostly due to re-running the route rather than being captured repeatedly.

SI3. Entering the nest does not improve re-running after being rewound in a section

SI3 Results

We compared ants' performance in Section A across the three treatments shown in Fig. SI3. ANOVAs showed significant differences across the three groups for both Scan rate ($F_{2,24} = 11.41$; $p < 0.001$) and Meander ($F_{2,24} = 11.39$; $p < 0.001$). Tukey's multi-comparison showed no difference between ZV ants tested immediately (Fig. SI3: 4th trial) and after entering the nest (Fig. SI3: Tested after nest ZV) (Scan rate: $p = 0.382$; Meander: $p = 0.996$). However, the ants that were tested as full vector after reaching the

feeder again (Fig. SI3: Tested after nest FV) showed significantly fewer scans ($ps < 0.004$) and lower Meander ($ps < 0.001$) than both ZV groups.

We also compared performance in Section A of the ants in condition 3c (tested as ZV ants after previously having entered their nest) to the ants in Condition 2a (rewound on A) and to the ants in Condition 2b (rewound on B) to calculate an evidence ratio of which Condition (2a or 2b) ants in Condition 3c resembled more. For both variables, ants in Condition 3c resembled the ants in Condition 2a more (Meander: evidence ratio 731.32; Scan rate: evidence ratio 3.25). These ratios, both above 1, mean overwhelming evidence in the case of meander and moderate evidence in the case of number of scans.

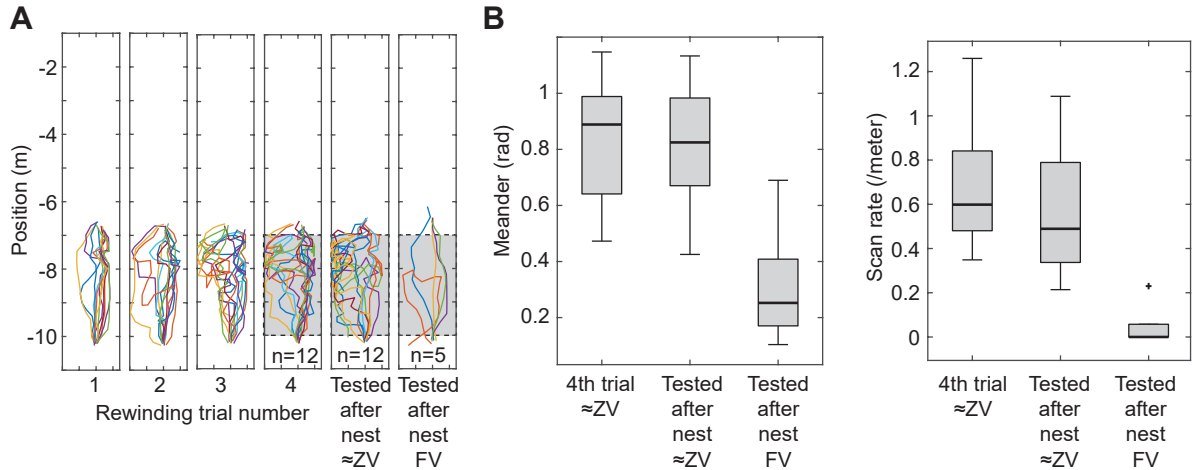


Fig. SI3. Entering the nest does not improve re-running after being rewound in a section. **A.** Paths of ants that re-ran Section A four times, then entered their nest, then were rewound in Section B four times on their next trip to reach a zero vector state, and then were tested on Section A either immediately (Condition 3c: Tested after nest =ZV), or on their next appearance at the feeder as full-vector ants (tested after nest FV). **B.** Meander (left) and Scan rate (right) in Section A (grey area in A) for the three treatments mentioned above. The box plots show the median (middle line in the box), the 25th and 75th quartiles (bottom and top of the box, respectively), and 1.5 times the

interquartile range (whiskers), as well as outliers (individuals beyond the range of the whiskers).

SI3 Discussion

It is clear that being in the nest did not alleviate the deleterious effects of repeated route running. However, the fact that FV ants' paths were much less affected than ZV ants' paths suggests that the path degradation observed in ZV ants resulted more from the suppression of appetitive route memories than learning new aversive visual memories that trigger turns. Indeed, given the suppression of appetitive route memories, ants should behave as in unfamiliar terrain, that is, follow their path integrator if the PI vector is large, and search around if in a ZV state, as we observed here. In the case of learning aversive memories, one would expect that the visual memories would dominate over the dictates of path integration, thus triggering turns in both ZV and FV ants. It should nonetheless be noted that the N is small, and that one of the FV ants did execute a U-turn, suggesting the existence as well of aversive memories, which may have arisen from being repeatedly captured at a specific location.

SI4. Set up and paths from Condition 4: comparing performance in an open vs. cluttered landscape

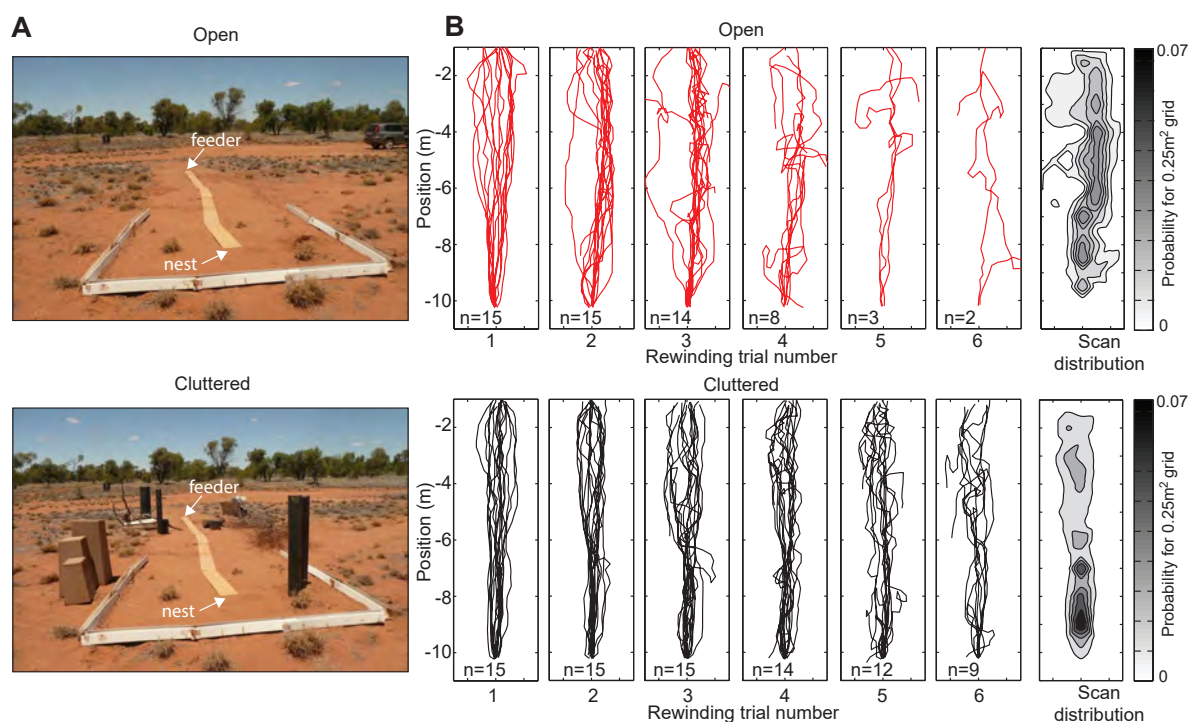


Figure SI4. Set up and paths from Condition 4: comparing performance in an open vs. cluttered landscape. **A.** Photos of the landscapes of the open condition (top) and the cluttered condition (bottom) of Nest 2. **B.** Paths (left 6 columns) and heat maps of scans (right column) of rewinding runs of ants in the open (top) and cluttered (bottom) conditions. In the heat map, the darker colours show a higher Scan rate. Quantitative analyses of these data are given in Fig. 3.